

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



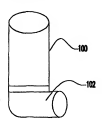
INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 7:		(11) International Publication Number:	WO 00/24362
A61K	A2	(43) International Publication Date:	4 May 2000 (04.05.00
(21) International Application Number: PCTUS (22) International Filing Date: 20 October 1999 (2 (30) Priority Data: 60105.850 27 October 1998 (27,10.58) 609(273,766 22 March 1999 (22.0.399) (71) Applicant: VIRGINIA COMMONWEALTH UNIV (USUS); 1101 E. Manshall Street, Richmond, V (US).	20.10.9 U TERSIT	BR. BY, CA. CH. CN. CR. C 9 ES, Fi, GB, GD, GE, GH, GM, KE, KG, KF, KR, KZ, LC, LK, MD, MG, MK, MN, MW, MX, SD, SE, SG, SI, SK, SI, TI, IS UZ, VN, YU, ZA, ZW, ARIPO MW, SD, SI, SZ, TZ, UG, ZW) BY, KG, KZ, MD, RU, TI, TM, CH, CY, DE, DK, ES, FI, FR, Y, LL, PT, SE), OAPI patent (BF,	U, CZ, DE, DK, DM, EF HR, HU, ID, II, IN, IS, JF LR, LS, LT, LU, LV, MA NO, NZ, PL, PT, RO, RL M, TR, TT, TZ, UA, UC patent (GH, GM, KE, LS, European patent (AM, AZ, European patent (AT, BE GB, GR, IE, IT, LU, MC BJ, CF, CG, CI, CM, GA
(72) Javentsers: PEART. Joanes, Apartment #332, 118t Wellesley Drive, Richmond, VA 23233 (US). BYS 18th June 1997. Per Per Very Richmond, VA 23233 (US). BYS 18th June 2325 (US). BYS 2525 (US). BYS 2	RON, P 129 (US ond, V ve, Ric	e- Without international search re upon receipt of that report.	port and to be republishe

(54) Title: Δ^9 TETRAHYDROCANNABINOL (Δ^9 THC) SOLUTION METERED DOSE INHALERS AND METHODS OF USE

(57) Abstract

The present invention provides therapeutic formulations for solutions of Δ^2 -termhydrocamabinol (Δ^2 THC) to be delivered by motered does inhalest. The formulations, which utilize non-CPC propellants, provide a stable acrosol-deliverable source of Δ^2 THC for the treatment of various medical conditions, such as mases and vormiting associated with chemotherapy; muscle spasticity; pairs, ancrexia associated with AIDS wasting syndrome; epilepsy; glaucoma, bronchial asthma; and mood disorders.



FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Paso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Gennany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		

Δ^{o} Tetrahydrocannabinol (Δ^{o} thc) solution metered dose inhalers and methods of use

DESCRIPTION

BACKGROUND OF THE INVENTION

Field of the Invention

The invention is generally related to the therapeutic use of Δ^{9} Tetrahydrocannabinol (Δ^{9} THC). In particular, the invention provides a metered dose inhaler (MDI) for the aerosol administration of Δ^{9} THC to patients suffering from nausea and vomiting associated with cancer chemotherapy, muscle spasticity, pain, anorexia associated with AIDS wasting syndrome, epilepsy, glaucoma, bronchial asthma. mood disorders, and the like.

Background Description

"Medical Marijuana" is a timely and controversial subject that is currently receiving widespread public attention. While marijuana is usually thought of as an illegal "recreational" drug, it also has a long history as a medicine. In 1997, the National Institutes of Health (NIH) released a review of the scientific data concerning potential therapeutic uses for marijuana. In that review, the NIH found that marijuana may indeed have beneficial medicinal effects and recommended that researchers develop alternative dosage forms for the drug, such as a "smoke free" inhaled delivery system (1). Table 1 summarizes the findings of several studies (references 2-18) that have documented therapeutically beneficial medicinal uses of the major active component of marijuana, Δ* tetrahydrocannabinol (Δ* THC).

TABLE 1. The Use of $\Delta^s THC$ for the Treatment of Assorted Clinical Conditions					
Condition and Number of Patients	Administration Route and Dose	Findings	Reference		
AIDS-associated anorexia and cachexia; 94 patients; 12 months	Oral placebo, 2.5 mg THC once or twice daily increasing to 20 mg daily	Long term THC treatment was well- tolerated; THC improved appetite and only tended to increase weight compared to controls	Beal et al., 1997		
AIDS-associated anorexia and cachexia; 139 patients; 42 days	Oral placebo or 2.5 mg THC twice daily	57% and 69% of vehicle and THC patients were evaluable for efficacy. Appetite increased 38% over baseline for THC group compared to only 8% for the placebo group. THC also decreased nausea. No significant changes were found between the groups for weight change.	Beal et al., 1995		
Nausea and emesis due to cancer chemotherapy; 36 patients who had experienced severe nausea and vomiting that was refractory to prochlorperazine or thiethylperazine	Oral THC, 15mg/m ²	Reduction in chemotherapy-induced nausea and vomiting in 64% of patients given THC compared to prochloperazine; side effects included dysphoria; authors recommend initial THC dose of 5 mg/m ²	McCabe et al., 1988		

TABLE 1. The Use of Δ^9 THC for the Treatment of Assorted Clinical Conditions				
Nausea and emesis due to cancer chemotherapy; 53 patients which were refractory to other antiemetics	Oral 5 or 15 mg/m ² THC four times per day	72% of patients exhibited a THC- induced partial or complete blockade of vomiting	Lucas and Laszlo, 1980	
Nausea and emesis due to cancer chemotherapy; 84 patients	Oral 10 mg/m² THC of prochloperazine	THC more effective than prochloperazine	Sallan et al., 1980	
Nausea and emesis due to cancer chemotherapy; 116 patients	Oral 15 mg THC, 10mg prochloperzine or placebo	Equal antiemetic effects between THC and prochlorperazine, effects of each greater than placebo; considerably more CNS side effects with THC than prochlorperazine	Frytak et al., 1979	
Nausea and emesis due to cancer chemotherapy; 15 patients	Oral placebo or 10 mg/m ² THC every 3 hours for a total of 5 doses, THC (17 mg) laced cigarettes of placebo were given if vomiting occurred	93% patients had a reduction in nausea and vomiting, 53% had an excellent response, 40% had a fair response; plasma THC levels 7,146.5 (mean ± SD) ng/ml. Side effects included sedation, tachycardia, few other side effects	Chang et al., 1979	
Pain due to advanced cancer; 10 patients	Oral placebo and 5, 10, 15 or 20mg THC	Pain relief, elevated mood, appetite stimulation, drowsiness, slurred speech, mental clouding	Noyes, et al, 1975	

TABLE 1. The Use Conditions	TABLE 1. The Use of $\Delta^9 THC$ for the Treatment of Assorted Clinical Conditions				
Pain due to advanced cancer; 34 patients	Placebo, 10 and 20mg THC, and 60 and 120 codeine	THC produced a similar degree of analgesia, with greater potency than codeine. THC CNS side effects included sedation, mental clouding, ataxia, and disorientation	Noyes et al. 1975		
Spasticity related to multiple schlerosis; 2 patients	Oral 10 or 15mg THC; rectal dose of 5 or 10mg THC	Improvement in passive mobility and walking ability	Brenneisen et al., 1996		
Spasticity related to multiple schlerosis; 13 patients	Oral 2.5 to 15mg THC once or twice daily or placebo	Significant subjective improvement in spasticity at 7.5mg THC and higher, no significant improvement in objective measurements	Ungerleider et al., 1987		
Spasticity related to multiple schlerosis; 8 patients, single blind	Oral 5 to 15mg THC	5 of 8 patients had mild subjective improvement in tremor. 2 of 8 patients had both objective and subjective improvement	Clifford, 1983		
Spasticity related to multiple schlerosis; 9 patients	Placebo, or 5 or 10mg THC	Decrease in spasticity compared to placebo treatment, minimal side effects	Petro and Ellenberger, 1981		
Spasticity and pain due to spinal cord injury; I patient	Oral placebo, THC (5 mg), or codeine (50 mg)	THC and codeine had analgesic effect compared to the placebo treatment. THC had a beneficial effect on spasticity whereas codeine did not	Maurer et al., 1990		

TABLE 1. The Use Conditions	TABLE 1. The Use of Δ^9 THC for the Treatment of Assorted Clinical Conditions				
Glaucoma, 6 patients	Oral placebo or 5, 10, 15 and 20 mg THC	Pain relief, elevated mood, appetite stimulation, drowsiness, slurred speech, mental clouding	Merritt et al, 1980		
Ten subjects with normal intra ocular pressure	Intravenous THC (0.022 or 0.044 mg/kg)	Decreased intra ocular pressure by a mean of 37%	Cooler and Gregg, 1977		
Nausea and emesis due to cancer chemotherapy; refractory to other antiemetics	Oral 10 mg/m² THC or placebo	In 20 courses of THC, 5 resulted in no vomiting, 9 resulted in a reduction of vomiting, 3 resulted in no decrease in vomiting, and 2 were unevaluable. THC was significantly better than placebo in decreasing vomiting.	Salian et al., 1975		

When marijuana is used illegally as a recreational psychoactive drug, the active ingredient Δ° THC is usually delivered to the lungs as an impure non-pharmaceutical aerosol in the form of marijuana smoke. Aerosolized Δ° THC in the inhaled smoke is absorbed within seconds and delivered to the brain efficiently. Table 2 and references 19-20 describe the pharmacokinetics of the administration of Δ° THC. As can be seen, inhalation is the preferred route of delivery for Δ° THC. When compared to oral delivery, inhalation provides a more rapid onset of pharmacological action and peak plasma levels. The effects achieved via inhalation are comparable to those achieved when the drug is administered intravenously, but inhalation is a much less invasive technique.

TABLE 2. Pharmacokinetics of ∆®THC Given Orally, Intravenously or by Smoking						
Route	Dose	% Dose in Plasma	Onset of Pharmacol ogical Action	Peak Plasma Levels	References	
Oral, sesame oil in gelatin capsules	2.5, 5, or 10 mg	10 to 20%	0.5 to 1 hour	12 0- 480 min	(PDR, 1995)	
Oral, in cookies	20 mg	4 to 12%	120-180 min	60-90 min	(Ohisson, et al.,	
Intra venous, bolus	5 mg	100%	10 min	3 min	(Ohlsson, et al., 1980)	
Smoking (THC lost to side stream smoke and pyrolysis	13 mg	8 to 24%	10 min	3 min	(Ohlsson, et al., 1980)	

Currently, the sources of Δ° THC for patients who could benefit from the drug are very limited. An oral form of Δ° THC (MARINOL) is marketed as a treatment for nausea and vomiting related to cancer chemotherapy, and as an appetite stimulant in patients suffering from AIDS wasting syndrome. In MARINOL, pharmaceutical grade Δ° THC is dissolved in sesame oil, encapsulated in gelatin capsules and delivered orally. However, when the drug is taken orally, the absorption is slower and more variable than when inhaled, with an onset of action between 30 minutes and 2 hours (Table 2). Alternatively, some cancer patients do manage to

obtain and smoke marijuana in order to alleviate such conditions as nausea and vomiting due to chemotherapy. This is, however, technically illegal and is thus obviously a less than ideal treatment protocol. There is no currently available pharmaceutically acceptable aerosol form of Δ° THC.

It would be advantageous to have available a form of pharmaceutical grade Δ° THC that could be administered as an aerosol. This would provide a means for rapid uptake of the drug without resorting to the illegal practice of smoking marijuana. Also, the potential adverse side effects encountered by smoking marijuana would be avoided. Further, an aerosol preparation of pharmaceutically pure Δ° THC could be administered in known, controlled dosages.

In 1976, Olsen et al. described a chlorofluorocarbon (CFC) propelled MDI formulation of Δ^9 THC (21). However, Δ^9 THC is known to deteriorate during storage, and the stability of Δ^9 THC in this formulation is suspect. In addition, the ethanol content in this formulation was so high (~23%) as to create an aerosol with droplets too large to be effectively inhaled (22). The Δ^9 THC CFC formulations were tested for use in treating asthma but were shown to be only moderately effective (23, 24). Moreover, CFC propellants have since been banned so that such a formulation in own useless. It would clearly be advantageous to develop a new aerosol formulation in which the Δ^9 THC is stable, the droplets are of a size that can be effectively inhaled, and which utilizes a non-CFC propellant.

SUMMARY OF THE INVENTION

It is an object of the present invention to provide a stable aerosol-dispensable pharmaceutical composition comprising a non-CFC propellant and a pharmaceutically effective concentration of Δ° THC. More particularly, it is an object of the present invention to provide a stable aerosol-dispensable pharmaceutical composition comprising a hydrofluoroalkane propellant, (for example, HFA 227 or HFA 134a) and Δ° THC. The propellant is present in the range of approximately 78 to 100% by

weight, and more particularly the propellant is present in the range of approximately 85 to 100% by weight. An organic solvent such as ethanol can be used to assist in solubilizing the Δ^{\bullet} THC in the propellant but is not required. If a solvent is used, preferably less than 20% by weight will be required, and most preferably less than 15% by weight will be required. The pharmaceutically effective concentration of Δ^{\bullet} THC is preferably in the range of 0.05 to 10% by weight, and most preferably in the range of 0.1 to 6% by weight. The pharmaceutical composition of the present invention can be used to treat a variety of medical conditions including nausea and vomiting associated with cancer chemotherapy, muscle spasticity, pain, anorexia associated with AIDS wasting syndrome, anorexia associated with cancer chemotherapy, epilepsy, glaucoma, bronchial asthma, mood disorders, migratine headaches.

DETAILED DESCRIPTION OF THE DRAWINGS

Figure 1. Δ^9 THC MDI characterization summary before and after storage at 40°C and 82% relative humidity (RH).

Figure 2. Generalized schematic drawings of a Δ9 THC MDI.

DETAILED DESCRIPTION OF A PREFERRED EMBODIMENT OF THE INVENTION

The instant invention provides a series of non-ozone depleting pressurized metered dose inhaler formulations of Δ^9 THC. In preferred embodiments of the invention, the formulations contain the pharmaceutically acceptable, non-ozone depleting hydrofluoroalkane propellants HFA 134a (1,1,1,2-tetrahydrofluoroethane) and HFA 227 (1,1,1,2,3,3,3-heptafluoropropane), or a mixture thereof.

When the propellant is a hydrofluoroalkane, it has been discovered that the propellant may be used with or without a solvent such as ethanol. Higher percentages of solvent generally allow higher levels of dissolution of Δ^0 THC. However, higher

percentages of solvent also cause droplet size to increase. In preferred embodiments of the invention, the range of propellant compositions, as shown in Table 3, may be from 100% propellant and 0% solvent to 85% propellant and 15% solvent. Within this range of percentages, pharmaceutically useful concentrations of Δ° THC can be achieved and droplet size is still small enough (<5.8 μ m) to provide excellent aerosol delivery of the drug. While these ratios reflect preferred embodiments of the invention, it will be recognized by those of skill in the art that the exact ratio of propellant to solvent (e.g. ethanol) may vary according to the desired final concentration of Δ° THC and droplet size. Any ratio of propellant to solvent that results in appropriate sized droplets and adequate dissolution of the Δ° THC may be used in the practice of this invention, and this will generally be in the range of from 100 to 80% propellant and 0 to 20% solvent. It is expected that a wide variety of solvents, such as ethanol, propanol, propylene glycol, glycerol, polyethylene glycol, etc. may be used in the preparation of formulations contemplated by this invention.

Those skilled in the art will also recognize that the "respirable dose" (or mass of Δ° THC in particles with aerodynamic diameters small enough to be delivered to and absorbed by the lungs) (Figure 1) may be increased by choosing MDI spray nozzles of different design and smaller orifice diameters. Respirable doses may also be increased by extending the mouthpiece of the MDI in such a way as to create an integral or separate aerosol spacer or reservoir attached to the mouthpiece of the MDI. This promotes an increase in droplet evaporation and hence in the percentage of the dose in smaller "respirable" particles or droplets. Generally, the optimal size of a respirable droplet is less than 10μ m in size.

TABLE 3. Apparent Solubility of Δ^9 THC in Ethanol/HFA Propellant Blends				
Formulation	Mass (g) of	Mass (g) of	Apparent	Comments
	Δ° THC in	Formulation	Solubility	
	Sample	Sampled	Mean (±SD)	-
Δ° THC in	0.000240	0.1071	0.224% w/w	Excess Δº THC added
100% HFA			(±0.063)	to propellant blend (in
134a				pressurized MDI).
		·		Solubility sample
				removed using puff
				absorber. n=5
Δ° THC in 5%	0.00144	0.0914	1.585% w/w	As above
Ethanol / 95%			(±0.321)	
HFA 134a				
Δ ⁹ THC in	0.00363	0.1036	3.511% w/w	As above
10% Ethanol /			(±0.249)	
90% HFA 134a				
Δ° THC in	0.00536	0.1098	4.883% w/w	As above
15% Ethanol /			(±0.224)	
85% HFA 134a				
Δ° THC in	0.00021	0.1451	0.147% w/w	As above
100% HFA 227			(±0.008)	
Δ° THC in 5%	0.00134	0.0979	1.339% w/w	As above
Ethanol / 95%			(±0.169)	
HFA 227				
Δ° THC in	0.00454	0.1267	3.240% w/w	As above
10% Ethanol/			(±0.161)	
90% HFA 227				
Δ° THC in	0.00623	0.1062	5.940% w/w	As above
15% Ethanol/			(±0.191)	
85% HFA 227			,	

A distinct advantage of the present formulations is that, surprisingly, the use of surface active agents or "surfactants" as valve lubricants and solubilizers is not necessary. This is in contrast to the invention of Purewal and Greenleaf (European Patent 0,372,777; reference #25) which provides HFA 134a/ethanol mixtures to produce stable formulations of pharmaceuticals in the presence of lipophilic surface active agents. Lipophilic surface active agents are incorporated in that invention in order to suspend undissolved material and to ensure adequate valve lubrication of the MDI. Without adequate valve lubrication, the useful life of the MDI and its ability to deliver an accurate dose of drug are severely attenuated. However, probably due to the inherent lubricity of the formulations of the present invention, the use of such surface active agents is unnecessary. This simplifies the composition and thus is an advantage with respect to cost and the elimination of potentially deleterious interactions between components of the formulations and the agents.

A major consideration in the formulation of any drug is its stability. Δ^{9} THC is known to deteriorate upon storage so that the effective concentration decreases and the purity is vitiated. The stability of the formulations of the present invention were tested according to accelerated storage testing protocols. The results are given in Figure 1 and Tables 4A and 4B. The formulations of the present invention were shown to be stable with respect to the release of aerosolized Δ^{9} THC in reproducible doses following accelerated storage testing. Apparently, the containment of Δ^{9} THC in solution in the non-aqueous formulations of the present invention is excellent with respect to chemical degradation, making possible the construction of a multidose inhaler with a good shelf life prognosis.

Further, lipophilic materials like Δ^o THC are generally known to partition into the elastomers of the valves in MDI formulations. (Δ^o THC is highly lipophilic as reflected in its octanol: water partition coefficient of 6000:1). Over time, this partitioning results in a decrease in the emitted or delivered dose of a lipophilic drug. Thus, this phemonemon also decreases the useful shelf-life of such preparations. However, the data presented in Figure 1 and Table 4 show that this is not the case

with the formulations of the present invention. The emitted or delivered doses were constant over the time period tested. This may be due to the somewhat surprising preference of Δ^9 THC for the formulation itself, rather than for the valve elastomers.

TABLE 4A. Formulation and aerosol characteristics of Δ° THC pressurized metered dose inhalers in ethanol/hydrofluoroalkane (HFA) propellant blends.

Inhaler	Fo	rmulation	Description	
	Δ' THC	Ethanol	Propellant	
1	0.13%	~5%	95% HFA 134a	3/98 Pale Yellow
				Solution
2	0.13%	~5%	95% HFA 227	3/98 Pale Yellow
				Solution
. 3	0.12%	~5%	95% HFA 134a	3/98 Pale Yellow
				Solution
4	0.18%	~5%	95% HFA 134a	3/98 Pale Yellow
				Solution
5	0.27%	~5%	95% HFA 227	3/98 Pale Yellow
				Solution
6	0.25%	~5%	95% HFA 134a	3/98 Pale Yellow
				Solution
7	0.57%	~5%	95% HFA 134a	3/98 Yellow Solution
8	0.58%	~5%	95% HFA 227	3/98 Yellow Solution
9	0.49%	~5%	95% HFA 134a	3/98 Yellow Solution
10	1.02%	~5%	95% HFA 134a	3/98 Yellow Solution
- 11	1.11%	~5%	95% HFA 227	3/98 Yellow Solution
12	0.97%	~5%	95% HFA 134a	3/98 Yellow Solution
SS* #1	1.07%	4.94%	94.0% HFA 134a	6/98 Yellow Solution
Initial				

SS* #1 after 28 days at 40°C/82% RH**	1.07%	4.94%	94.0% HFA 134a	7/98 Yellow Solution
SS* #2 after 21 days at 40°C/82% RH**	1.00%	5.01%	95% HFA 134a	7/98 Yellow Solution
SS* #3 Modified Actuator**	1.02%	5.15%	93.8% HFA 134a	10/98 Yellow Solution

^a Mean (Standard Deviation) of five determinations.

TABLE 4B. Formulation and aerosol characteristics of Δ^9 THC pressurized metered dose inhalers in ethanol/hydrofluoroalkane (HFA) propellant blends.

Inhaler	Aero	sol Characterization	
	Metered Dose (mg) ^a	Emitted Dose (mg) ^a	Fine Particle Dose (mg)
11	1.72 (0.25)	1.32 (0.17)	ND
12	0.94 (0.23)	0.97 (0.10)	0.38 (0.02)

 $^{^{\}text{b}}$ Mass of Δ^{9} THC aerosol particles <5.8 μm aerodynamic diameter

^{*}SS: Stability Sample

^{**} RH: relative humidity

^{***} Aproximate spray nozzle diameter = 0.2 mm.

SS* #1 Initial	1.10 (0.07)	0.90 (0.03)	0.22 (0.03)
SS* #1 after 28 days at 40°C/82% RH**	1.06 (0.03)	0.92 (0.04)	0.23 (0.02)
SS* #2 after 21 days at 40°C/82% RH**	1.02 (0.05)	0.90 (0.05)	0.21 (0.02)
SS* #3 Modified Actuator***	ND .	ND	0.40 (n=1)

a Mean (Standard Deviation) of five determinations.

** RH: relative humidity

ND: not determined

The final concentration of Δ^o THC in a given formulation may be varied by adjusting the ratio of propellant to solvent and thus the solubility of the Δ^o THC. Higher percentages of solvent (e.g. ethanol) generally allow a higher amount of Δ^o THC to be dissolved. For example, in preferred embodiments of the invention, the apparent solubility of Δ^o THC ranged from 0.147% w/w to 5.94% w/w as the propellant composition varied from 100% HFA 227 to 85% HFA 227 and 15% ethanol. Thus, the dose of Δ^o THC in a given metered volume may be selected by changing the formulation.

Further, as stated above, the "fine particle dose" or "respirable dose" of a drug dispensed with an MDI is a function of the spray nozzle diameter. In Figure 1 and Tables 4A and 4B, the spray nozzle diameter is 0.4mm. The "fine particle dose" or "respirable dose" of the formulations of the present invention was shown to be

 $^{^{\}text{b}}$ Mass of Δ^9 THC aerosol particles with <5.8 μm aerodynamic diameter

^{*}SS: Stability Sample **

^{***} Approximate spray nozzle diameter = 0.2 mm

unaffected by storage.

The Δ° THC of the present invention is pharmaceutically pure. That is, its form is the nonionized resinous drug substance (6aR-trans)-6a,7,8,10a-tetrahydro-6,6,9trimethyl-3-pentyl-6H-dibenzo[b,d]-pyran-1-ol. Although its preferred embodiment in this invention is not a sait or ester, it will be readily understood by those of skill in the art that other appropriate forms of Δ° THC may be synthesized (e.g. esters and salts) and thus used in the practice of this invention.

The desired final concentration of Δ° THC in a patient's serum will vary from patient to patient depending on, for example, the nature and severity of the condition being treated, and the patient's overall condition, weight, gender and response to the drug, etc. But the desired range will generally be 10-100ng/ml at 15 minutes following inhalation. The level of Δ° THC in a patient's serum can be readily and reliably monitored by gas chromatography/mass spectrophotometry (GC/MS).

The exact treatment protocol to be used may vary from patient to patient depending on the circumstances. For example, in a preferred embodiment of the invention, a patient receiving chemotherapy may have one dose of Δ^9 THC prescribed via inhalation, to be administered 15 minutes before chemotherapy and 4-8 times daily following chemotherapy. In another preferred embodiment, a patient suffering from anorexia associated with AIDS wasting syndrome may have Δ^9 THC by inhalation prescribed 3-5 times daily, 30 minutes before each meal or snack. In other preferred embodiments, a patient suffering form cancer pain, or spaticity related to either multiple sclerosis or spinal cord injury may have Δ^9 THC by inhalation prescribed 3-6 times daily. Those skilled in the art will readily recognize that the treatment protocol may be crafted so as to address the particular needs of each individual patient on a case by case basis.

 Δ^{9} THC may be used alone or in combination with other medications. Those skilled in the art will readily recognize that, for example, in the case of AIDS wasting syndrome, the patient will likely also be taking drugs that combat the AIDS virus. Similarly, those skilled in the art will readily recognize that patients receiving

chemotherapy for cancer may also receive other antiemetics, and cancer patients seeking to relieve pain are likely to receive opioids as well as nonsteroidal antiinflammatory agents.

The containers for the formulations of the instant invention may be any that are suitable for the efficacious delivery of aerosol inhalants. Several containers and their method of usage are known to those of skill in the art. For example, MDIs can be used with various dose metering chambers, various plastic actuators and mouthpieces, and various aerosol holding chambers (e.g. spacer and reservoir devices), so that appropriate doses of Δ^9 THC cach and deposit in the lung and are thereafter absorbed into the bloodstream. In addition, a lock mechanism such as that shown in U.S. Patent 5,284,133 to Burns and Marshak, which is herein incorporated by reference, can be used to prevent overdose or unauthorized consumption of Δ^9 THC. Figure 2 provides a generalized drawing of an MDI containing the composition of this invention and provides the advantage of delivering metered quantities of Δ^9 THC on a repetitive basis. The MDI includes a container 100 for holding the composition and a valve delivery mechanism 102 for delivery of aerosolized Δ^9 THC.

While the invention has been described in terms of its preferred embodiments, those skilled in the art will recognize that the invention can be practiced with modification within the spirit and scope of the appended claims.

REFERENCES

- Workshop on the medical utility of marijuana. National Institutes of Health, August 1997.
- Beal, J.A., Olson, R., Lefkowitz, L., Laubenstein, L., Bellman, P., Yangco, B., Morales, J.O., Murphy, R., Powderly, W., Plasse, T.F., Mosdell, K.W. and Shepard, K.W. (1997) Long-term efficacy and safety of dronabinol for acquired immunodeficiency syndrome-associated anorexia. J. Pain. Symptom Manage. 14:7-14.

 Beal, J.A., Olson, R., Laubenstein, L., Morales, J.O., Bellman, B., Yangco, B., Lefkowitz, L., Plasse, T.F. and Shepard, K.V. (1995) Dronabinol as a treatment for anorexia associated with weight loss in patients with AIDS J. Pain. Symptom Manage. 10:89-97.

- McCabe, M., Smith, F.P., MacDonald, J.S., Wooley, P.V., Goldberg, D. and Schein, P.S. (1988) Efficacy of tetrahydrocannabinol in patients refractory to standard antiemetic therapy. *Invest. New Drugs* 6:243-246.
- 5. Lucas, V.S. and Laszlo, J. (1980) Δ^9 -THC for refractory vomiting induced by cancer chemotherapy. *JAMA* 243:1241-1243.
- Sallan, S.E., Cronin, C., Zelen, M.and Zinberg, N.E. (1980) Antiemetics in patients receiving chemotherapy for cancer: a randomized comparision of Δ⁵ THC and prochlorperazine. N. Engl. J. Med. 302:135-138.
- 7. Frytak, S., Moertel, C.G., O'Fallon, J.R., Rubin, J., Creagan, E.T., O'Connell, M.J., Schutt, A.J. and Schwartau, N.W. (1979) Delta-9-tetrahydrocannabinol as an antiemetic for patients receiving cancer chemotherapy: a comparison with prochlorperazine and a placebo. Ann. Inter. Med. 91:825-830.
- Chang, A.E., Shiling, D.J., Stillman, R.C., Goldgerg, N.H., Seipp, C.A., Barofdky, I., Simon, R.M. and Rosenberg SA (1979) \(\Delta^{\text{o}}\) THC as an antiemitic in cancer patients receiving high-dose methotrexate. Ann. Internal. Med. 91:819-824.
- Sallan, S.E., Zinberg, N.E. and Frei, I.E. (1975) Antiemetic effect of Δ^o THC in patients receiving cancer chemotherapy. New Engl. J. Med. 293:795-797.
- 10. Noyes, J.R., Brunk, S.F., Baram, D.A. and Canter, A. (1975) The analgesic

properties of Δ9 THC and codeine. J. Clin. Pharmacol. 15:139-143.

- 11. Noyes, R., Jr., Brunk, S.F., Baram, D.A. and Canter, A. (1975) Analgesic effect of Δ^9 -tetrahydrocannabinol. J. Clin. Pharmacol. 15:139-143.
- 12. Brenneisen, R., Egli, A., Elosohlly, M.A., Henn, V. and Spiess, Y. (1996) The effect of orally and rectally administered Δ^0 THC on spasticity: a pilot study with 2 patients. Int. J. Clin. J. Pharmocol. Ther. 34:446-452.
- Ungerleider, J.T., Andyrsiak, T.F.L., Ellison, G.W. and Myers, L.W. (1987) Δ⁹
 THC in the treatment of spasticity associated with multiple sclerosis. Adv. Alcohol Subst. Abuse 7:39-50.
- Clifford, D.B. (1983) Tetrahydrocannabinol for tremor in multiple sclerosis. Ann. Neurol. 13:669-171.
- Petro, D.J. and Ellenberger, C. (1981) Treatment of human spasticity with delta 9 -tetrahydrocannabinol. J. Clin. Pharmacol. 21:413S-416S.
- 16. Maurer, M., Henn, V., Dittrich, A. and Hofman, A. (1990) Delta 9tetrahydrocannabinol shows antispastic and analgesic effects in a single case doubleblind trial. Eur. Arch. Psychiatry Neurol. Sci. 240:1-4.
- Merritt, J., Crawford, W., Alexander, P., Anduze, A. and Gelbart, S. (1980)
 Effects of marihuana on intra ocular and blood pressure in glaucoma. Opht. 87:222-228.
- 18. Cooler, P. and Gregg, J.M. (1977) Effect of delta 9- Δ° THC on intra ocular pressure in humans. South. Med. J. 70:951-954.

19. PDR (1995) Physician's Desk Reference (49) Montvalek, New Jersey: Medical Economics Data Production Co., pp.2787.

- Ohlsson, A., Lindgren, J.E., Wahlen, A., Agurall, S., Hollister, L.E. and Gillespie, H.K. (1980) Plasma Δ⁹ THC concentrations and effects after oral and intravenous administration and smoking. Clin. Pharmacol. Ther. 28:409-416.
- 21. Olsen, J.L., Lodge, J.W., Shapiro, B.J. and Tashkin, D.P. (1976) An inhalation aerosol of Δ° tetrahydrocannabinol. *J. Pharmacy and Pharmacol.* 28:86.
- Dalby, R.N. and Byron, P.R. (1988) Comparison of output particle size distributions from pressurized aerosols formulated as solutions or suspensions. *Pharm. Res.* 5:36-39.
- Tashkin, D.P., Reiss, S., Shapiro, B.J., Calvarese, B., Olsen, J.L. and Lidgek, J.W. (1977) Bronchial effects of aerosolized
 ² tetrahydrocannabinol in healthy and asthmatic subjects. Amer. Rev. of Resp. Disease. 115:57-65.
- 24. Williams, S.J., Hartley, J.P.R. and Graham, J.D.P. (1976) Bronchodilator effect of delta-9-THC administered by aerosol to asthmatic patients. *Thorax*, 31:720-723.
- 25. European Patent 0,372,777 (Riker Laboratories). Medicinal aerosol formulations.

CLAIMS

	We claim:
1	1. An aerosol-dispensable pharmaceutical composition, comprising:
2	a hydrofluoroalkane propellant; and

3

4

5

1

2

3

1

2

2

2

1

3

 $\Delta^9\text{-}$ tetrahydrocannabinol present in a pharmaceutically acceptable form and at a

pharmaceutically effective concentration dissolved in said hydrofluoroalkane propellant.

- 2. The aerosol dispensable pharmaceutical composition of claim 1 wherein said Δ^{9} tetrahydrocannabinol is dissolved in said hydrofluoroalkane propellant without the use of a solubilizing agent selected from the group consisting of solvents and surfactants.
- The aerosol-dispensable pharmaceutical composition of claim 1 wherein said hydrofluoroalkane is selected from the group consisting of: HFA 134a and HFA 227.
- The aerosol-dispensable pharmaceutical composition of claim 3 wherein said hydrofluoroalkane is present in an amount in excess of 85% by weight.
- The aerosol-dispensable pharmaceutical composition of claim 1 further comprising an organic solvent.
- The aerosol-dispensable pharmaceutical composition of claim 5 wherein said organic solvent is ethanol present in an amount ranging up to 15% by weight.
- 7. The aerosol-dispensable pharmaceutical composition of claim 1 wherein said pharmaceutically effective concentration of Δ^0 -tetrahydrocannabinol ranges from 0.05 to 10% by weight.

1	
2	8. The aerosol-dispensable pharmaceutical composition of claim 1 wherein said
3	pharmaceutically effective concentration of \$\Delta^9\$-tetrahydrocannabinol ranges from 0.1
4	to 6% by weight.
1	9. The aerosol-dispensable pharmaceutical compositon of claim 1 wherein Δ^9 -
2	tetrahydrocannabinol is present in pure form.
1	10. A solvent free pharmaceutical composition consisting essentially of
2	1,1,1,2,3,3,3-hydrofluoropropane (HFA 227) and Δ^2 - tetrahydrocannabinol.
ı	11. A solvent free pharmaceutical composition consisting essentially of
2	1,1,1,2-hydrofluroethane (HFA 134a) and $\Delta^{\circ}\text{-}$ tetrahydrocannabinol.
1	12. A method of treating a patient in need thereof with an aerosolized
.2	pharmaceutically acceptable form of Δ^9 - tetrahydrocannabinol, comprising the step of:
3	administering an aerosolized dose of a pharmaceutically acceptable form of
4	Δ^9 -tetrahydrocannabinol as respirable droplets to a patient's lung from a composition
5	comprised of a hydrofluoroalkane propellant and said pharmaceutically acceptable
6	form of Δ^{9} - tetrahydrocannabinol.
1	13. The method of claim 12 wherein said composition comprises a
2	pharmaceutically acceptable solvent.
3	14. The method of claim 12 wherein said composition is solvent-free.
4	15. The method of claim 12 wherein said aerosolized dose is sufficient to
5	reduce nausea.

1	16. The method of claim 12 wherein said aerosolized dose is sufficient to
2	reduce vomiting.
1	17. The method of claim 12 wherein said aerosolized dose is sufficient to
2	reduce pain.
1	18. The method of claim 12 wherein said aerosolized dose is sufficient to
2	relieve muscle spasticity.
1	10. The marked of alain 10 art and a state of the state o
2	19. The method of claim 12 wherein said aerosolized dose is sufficient to relieve migraine headaches.
2	reneve inigrame neadaches,
1	20. The method of claim 12 wherein said aerosolized dose is sufficient to
2	relieve movement disorders.
1	21. The method of claim 12 wherein said aerosolized dose is sufficient to
2	increase appetite in patients suffering from cachexia.
1	22. The method of claim 12 wherein said pharmaceutically acceptable form of
2	Δ^9 - tetrahydrocannabinol is pure Δ^9 - tetrahydrocannabinol and said hydrofluoroalkane
3	is selected from the group consisting of HFA 134a and HFA 227.
4	23. A metered dose inhaler, comprising,
5	a housing;
6	a metering valve connected to said housing; and.
7	an aerosol-dispensable pharmaceutical composition which includes a
8	hydrofluoroalkane propellant and Δ^{9} - tetrahydrocannabinol present in a
9	pharmaceutically effective concentration dissolved in said hydrofluoroalkane
0	propellant.
•	propermit

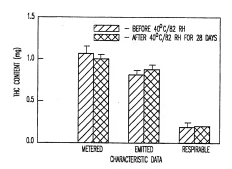


FIG.1

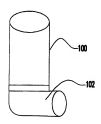


FIG.2